Liaison Division, provides various personnel management services to a number of diverse Presidential commissions, committees, boards and other agencies through reimbursable administrative support agreements. This notice is proceeded on behalf of the client agencies, and it supersedes all other notices in the **Federal Register** on this subject.

Because of their small size, a Performance Review Board register has been established in which SES members from the client agencies participate. The Board is composed of SES members from various agencies. From this register of names, the head of each client agency will appoint executives to a specific board to serve a particular client agency.

The members whose names appear on the Performance Review Board standing roster to sere client agencies are:

Administrative Conference of the U.S. Gary J. Edles, General Counsel Jeffrey S. Lubbers, Research Director

Barry M. Goldwater Scholarship and Excellence in Education Foundation Gerald J. Smith, Executive Secretary

Board of International Broadcasting

Richard McBride, Executive Director John A. Lindburg, General Counsel Patricia H. Schlueter, Director of Financial and Congressional Affairs Bria T. Conniff, Inspector General

Committee for Purchase From People Who Are Blind or Severely Disabled

Beverly L. Milkman, Executive Director

Defense Nuclear Facilities Safety Board

Kenneth M. Pusateri, General Manager Joseph R. Neubeiser, Deputy General Manager

Robert M. Anderson, General Counsel Richard A. Azzaro, Deputy General Counsel for Policy and Litigation

George W. Cunningham, General Engineer Joyce P. Davis, Chief, Health Physics Branch Wallace R. Kornack, Assistant Director for Engineering

Steven L. Krahn, Assistant Director for Weapon Programs

Lester A. Ettlinger, Assistant Director for Standards

Harry S Truman Scholarship Foundation Louis H. Blair, Executive Secretary

Japan-United States Friendship Commission

Office of Navajo and Hopi Indian Relocation Christopher J. Bavasi, Executive Director Michael J. McAlister, Deputy Executive Director

Artic Research Commission
Garrett W. Brass, Executive Director
National Mediation Board
Ronald M. Etters, General Counsel

Eric J. Gangloff, Executive Director

Dated: July 18, 1995.

Calvin R. Snowden,

Director.

[FR Doc. 95-18486 Filed 7-26-95; 8:45 am] BILLING CODE 6820-34-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

CDC Recommendations for Civilian Communities Near Chemical Weapons Depots: Guidelines for Medical Preparedness; Correction

A notice was published in the **Federal Register** on June 27, 1995 (60 FR 33308), entitled, "CDC Recommendations for Civilian Communities Near Chemical Weapons Depots: Guidelines for Medical Preparedness." This notice is corrected as follows:

On page 33309, first column, line 6 of the second paragraph, change "falls" to "fall"; and in the second column, line 9, under the heading: 1. Personal Protective Equipment (PPE), change "have" to "has" and in line 11, change "portable" to "powered." On page 33311, second column, line 8, under the heading: 5. Personal Protective Equipment (PPE), change "have" to "has."

Dated: July 21, 1995.

Joseph R. Carter,

Acting Associate Director for Management and Operations, Centers for Disease Control and Prevention (CDC).

[FR Doc. 95-18433 Filed 7-26-95; 8:45 am] BILLING CODE 4163-18-P

Food and Drug Administration

Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). This notice also summarizes the procedures for the meeting and methods by which interested persons may participate in open public hearings before FDA's advisory committees.

FDA has established an Advisory Committee Information Hotline (the hotline) using a voice-mail telephone system. The hotline provides the public with access to the most current information on FDA advisory committee meetings. The advisory committee hotline, which will disseminate current information and information updates, can be accessed by dialing 1–800–741–8138 or 301–443–0572. Each advisory committee is assigned a 5-digit number. This 5-digit number will appear in each individual notice of meeting. The hotline will enable the public to obtain information about a particular advisory committee by using the committee's 5-digit number. Information in the hotline is preliminary and may change before a meeting is actually held. The hotline will be updated when such changes are made.

MEETING: The following advisory committee meeting is announced:

Subcommittee Meeting of the National Task Force on AIDS Drug Development on Drug Development Issues

Date, time, and place. September 13 and 14, 1995, 8:30 a.m., Gaithersburg Hilton Hotel, Grand Ballroom, 629 Perry Pkwy., Gaithersburg, MD. A limited number of overnight accommodations have been reserved at the Gaithersburg Hilton Hotel. Attendees requiring overnight accommodations may contact the hotel at 301–977–8900 and reference the task force. Interested persons are encouraged to register early because space is limited.

Type of meeting and contact person. Open subcommittee discussion, September 13, 1995, 8:30 a.m. to 11:45 a.m.; open public hearing, 11:45 a.m. to 12:15 p.m., unless public participation does not last that long; open subcommittee discussion, 12:15 p.m. to 4:45 p.m.; open subcommittee discussion, September 14, 1995, 8:30 a.m. to 11:45 a.m.; open public hearing, 11:45 a.m. to 12:15 p.m., unless public participation does not last that long; Nancy L. Stanisic, Office of AIDS and Special Health Issues (HF-12), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-443-0104, or FDA Advisory Committee Information Hotline, 1-800-741-8138 (301–443–0572 in the Washington, DC area), National Task Force on AIDS Drug Development, code 12602.

General functions of the task force. The task force shall identify any barriers and provide creative options for the rapid development and evaluation of treatments for human immunodeficiency virus (HIV) infection and its sequelae. It also advises on issues related to such barriers, and provides options for the elimination of these barriers.

Open subcommittee discussion. The subcommittee will review, discuss, and clarify issues concerning the

investigational new drug applications and submissions to the National Institutes of Health's Recombinant DNA Advisory Committee as it relates to gene therapy. Representatives from FDA's Center for Biologics Evaluation and Research and the National Institutes of Health's Office of Recombinant DNA Activities will make presentations and answer questions concerning the application process.

Agenda—Open public hearing. Interested persons may present data, information, or views, orally or in writing, on issues pending before the task force. Those desiring to make formal presentations should notify the contact person before September 6, 1995, and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time required to make their comments.

FDA public advisory committee meetings may have as many as four separable portions: (1) An open public hearing, (2) an open committee discussion, (3) a closed presentation of data, and (4) a closed committee deliberation. Every advisory committee meeting shall have an open public hearing portion. Whether or not it also includes any of the other three portions will depend upon the specific meeting involved. There are no closed portions for the meetings announced in this notice. The dates and times reserved for the open portions of each committee meeting are listed above.

The open public hearing portion of each meeting shall be at least 1 hour long unless public participation does not last that long. It is emphasized, however, that the 1 hour time limit for an open public hearing represents a minimum rather than a maximum time for public participation, and an open public hearing may last for whatever longer period the committee chairperson determines will facilitate the committee's work.

Public hearings are subject to FDA's guideline (subpart C of 21 CFR part 10) concerning the policy and procedures for electronic media coverage of FDA's public administrative proceedings, including hearings before public advisory committees under 21 CFR part 14. Under 21 CFR 10.205, representatives of the electronic media may be permitted, subject to certain limitations, to videotape, film, or otherwise record FDA's public administrative proceedings, including presentations by participants.

Meetings of advisory committees shall be conducted, insofar as is practical, in accordance with the agenda published in this **Federal Register** notice. Changes in the agenda will be announced at the beginning of the open portion of a meeting.

Any interested person who wishes to be assured of the right to make an oral presentation at the open public hearing portion of a meeting shall inform the contact person listed above, either orally or in writing, prior to the meeting. Any person attending the hearing who does not in advance of the meeting request an opportunity to speak will be allowed to make an oral presentation at the hearing's conclusion, if time permits, at the chairperson's discretion.

The agenda, the questions to be addressed by the committee, and a current list of committee members will be available at the meeting location on the day of the meeting.

Transcripts of the open portion of the meeting may be requested in writing from the Freedom of Information Office (HFI-35), Food and Drug Administration, rm. 12A–16, 5600 Fishers Lane, Rockville, MD 20857, approximately 15 working days after the meeting, at a cost of 10 cents per page. The transcript may be viewed at the Dockets Management Branch (HFA-305), Food and Drug Administration, rm. 1-23, 12420 Parklawn Dr., Rockville, MD 20857, approximately 15 working days after the meeting, between the hours of 9 a.m. and 4 p.m., Monday through Friday. Summary minutes of the open portion of the meeting may be requested in writing from the Freedom of Information Office (address above) beginning approximately 90 days after the meeting.

This notice is issued under section 10(a)(1) and (2) of the Federal Advisory Committee Act (5 U.S.C. app. 2), and FDA's regulations (21 CFR part 14) on advisory committees.

Dated: July 19, 1995.

Linda A. Suydam,

Interim Deputy Commissioner for Operations. [FR Doc. 95–18503 Filed 7–26–95; 8:45 am] BILLING CODE 4160–01–F

Health Care Financing Administration

Privacy Act of 1974; Report of New System

AGENCY: Department of Health and Human Services (HHS), Health Care Financing Administration (HCFA). **ACTION:** Notice of new system of records.

SUMMARY: In accordance with the requirements of the Privacy Act of 1974, we are proposing to establish a new system of records, "Evaluation of, and

External Quality Assurance for, the Community Nursing Organization (CNO) Demonstration," HHS/HCFA/ORD No. 360–94–30500, 30501. We have provided background information about the proposed system in the "Supplementary Information" section below. Although the Privacy Act requires only that the "routine uses" portion of the system be published for comment, HCFA invites comments on all portions of this notice.

DATES: HCFA filed a new system report with the Chairman of the Committee on Government Operations of the House of Representatives, the Chairman of the Committee on Governmental Affairs of the Senate, and the Administrator, Office of Information and Regulatory Affairs, Office of Management and Budget (OMB), on July 21, 1995. To ensure that all parties have adequate time in which to comment, the new system of records, including routine uses, will become effective 40 days from the publication of this notice or from the date the report was submitted to OMB and the Congress, whichever is later, unless HCFA receives comments which require alterations to this notice. **ADDRESSES:** The public should address comments to Richard DeMeo, HCFA Privacy Act Officer, Office of the Associate Administrator for External Affairs, HCFA, Room C2-01-11, 7500 Security Boulevard, Baltimore,

FOR FURTHER INFORMATION CONTACT: Melissa McNiff, Project Officer for the evaluation of the Community Nursing Organization Demonstration and the

received will be available for inspection

Maryland 21244–1850. Comments

at this location.

Organization Demonstration and the External Quality Assurance Program, Office of Research and Demonstrations, HCFA, Room C3–21–06, 7500 Security Boulevard, Baltimore, Maryland 21244–1850, Telephone 410–786–8494.

SUPPLEMENTARY INFORMATION: HCFA proposes to initiate a new system of records, collecting data under the authority of section 4079 of Pub. L. 100-203, the Omnibus Budget Reconciliation Act of 1987. The purpose of this system is to provide data necessary to test the operational feasibility of the CNO and examine whether the combination of capitated payment and nurse-case management will promote timely and appropriate use of community nursing and ambulatory care services and reduce the use of costly acute care services. It will further determine the effect of membership in a CNO on a typical beneficiary's use of health services covered under the CNO package and on services such as physician and inpatient hospital care